1 2	<u>CLAIMS</u>
3 4	What is claimed is:
5	Claim 1. A method for diagnosing congestive heart
6	failure (CHF) in a subject, comprising the steps of:
7	A) contacting a monoclonal antibody specific for a
8	glycophorin antigen with a biological fluid obtained from
9	said subject under conditions such that an antibody-antigen
10	binding complex forms between said monoclonal antibody and
11	said glycophorin antigen present in said biological fluid;
12	and
13	B) detecting said antibody-antigen binding complex
14	wherein the presence of said antibody-antigen binding complex
15	is diagnostic for congestive heart failure (CHF).
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17	Claim 2. The method in accordance with claim 1, wherein
18	said biological fluid is selected from the group consisting
19	of blood, blood products, urine, saliva, cerebrospinal fluid
20	and lymphatic fluid.
21	
22	Claim 3. The method in accordance with claim 1, wherein
23	said monoclonal antibody is 3F4 and recognizes amino acid
24	residues 5-25 of SEQ ID NO:2 and SEQ ID NO:4.
25	

1	Claim 4. The method in accordance with claim 1, wherein
2	said monoclonal antibody is 6G4 and recognizes amino acid
3	residues 39-45 of SEQ ID NO:2.
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5	Claim 5. The method in accordance with claim 1, wherein
6	said monoclonal antibody is 5F4 and recognizes amino acid
7	residues 107-119 of SEQ ID NO:2.
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9	Claim 6. The method in accordance with claim 1, wherein
10	said glycophorin antigen is a truncated glycophorin.
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12	Claim 7. The method in accordance with claim 1, wherein
13	said detecting comprises the steps of:
14	A) contacting said antibody-antigen binding complex with
15	a polyclonal antibody corresponding to said glycophorin
16	antigen under conditions such that a complex forms between
17	said glycophorin antigen and said polyclonal antibody;
18	B)attaching a label to a polyclonal antibody
19	corresponding to the polyclonal antibody of step A;
20	C) contacting the complex formed in step A with the
21	labeled polyclonal antibody formed in step B under conditions
22	such that a complex forms between said labeled polyclonal
23	antibody and said polyclonal antibody of step A; and
24	C) detecting the label on said labeled polyclonal
25	antibody.

Claim 8. The method in accordance with claim 7, wherein the label on said labeled polyclonal antibody comprises a signal generating substance. Claim 9. The method in accordance with claim 8, wherein said signal generating substance is peroxidase.